NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 20

STEREO ATTRIBUTES: NONE

=> s 16 ful FULL SEARCH INITIATED 15:05:58 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 1283 TO ITERATE

100.0% PROCESSED 1283 ITERATIONS SEARCH TIME: 00.00.01

194 ANSWERS

L7 194 SEA SSS FUL L6

=> fil caplus COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION 361.07 FULL ESTIMATED COST 360.86

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=> s 17
L8
              5 L7
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=> d bib abs 1-5

- ANSWER 1 OF 5 CAPLUS COPYRIGHT 2008 ACS on STN
- 2005:34761 CAPLUS AN
- 142:113915 DN
- TI Preparation of heterocyclic substituted 4-(aminomethyl)piperidine benzamides as 5-HT4 antagonists
- TN Bosmans, Jean-Paul Rene Marie Andre; Gijsen, Henricus Jacobus Maria; Mevellec, Laurence Anne
- Janssen Pharmaceutica N. V., Belg. PA
- PCT Int. Appl., 46 pp. SO
- CODEN: PIXXD2 DT Patent

LA FAN.	CNT							D3 MD				T 0 3 M	TO11							
	PATENT NO.						KIND DATE				APPLICATION NO.									
PI		2005														2	0040	610		
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			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,		
								ID,												
								LV,												
								PL,												
								TZ,												
		RW:						MW,												
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								GR,												
						Br,	ы,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MK,	NE,		
	2.11	2004		TD,		3.1		2006	0112		7.11 O	004	25.41	0.0		2	0040	610		
										AU 2004-254190										
										CA 2004-2528642 EP 2004-739776										
	EF							ES,												
		14.						RO,											uп	
	.TP	2006																	111/	
		2007																		
PRAT		2003									-	000	5001	00		_	0001			
2.411		2003																		

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB The title compds. I [R1R2 = O(CH2)nO, n = 1-4; R1R2 = O(CH2)n, m = 2-5; R3 = H, halide, C1-C6-alkyl, C1-C6-alkyl, C1-C6-alkyl, C1-C6-alkyl, C1-C6-alkyl, C1-C6-alkyl, C1-C6-alkyl, NH2, mono or di(C1-C6-alkyl)amino; R5 = H, C1-C6-alkyl, OR5 = 3- or 4-position; L = C1-C12-alkanediyl-R6, C1-C12-alkanediyl-X-R7; R6, R7 = heterocycle or heterocycle substituted with halide, OH, C1-C6-alkyl; heterocycle = morpholine, tetracole, pyrazole, isoxazole, isoxiniazole, oxazolyl, thiazole, pyran, 2,4-dioxoimidazolidine] were prepared and tested as 5-HT4 antagonists. For example, reacting (chloropropyl)trityltetrazole II with trans-hydroxypiperidine derivative III gave alkylated piperidine IV (R = CPh3) which was deprotected to give IV (R = H).

RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L8 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2008 ACS on STN
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AN 2005:34760 CAPLUS

DN 142:134469

TI A preparation of 5HT4-antagonistic N-(piperidin-4-ylmethyl)-benzamide derivatives

IN Bosmans, Jean-Paul Rene Marie Andre; Gijsen, Henricus Jacobus Maria; Mevellec, Laurence Anne

PA Janssen Pharmaceutica N. V., Belg.

SO PCT Int. Appl., 63 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

E MIN.	PA	TENT												NO.			ATE		
PI		2005															0040	610	
		W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	
			LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NA,	NI,	
			NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	
			ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	zw	
		RW:						MW,											
								RU,											
								GR,											
						BF,	ΒJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,	
				TD,															
										AU 2004-254193 CA 2004-2528656					20040610				
								20060329			EP 2	004-	-736513			2	200406	610	
	EP	1638																	
		R:						ES,											
								RO,											
		P 2006527715																	
		3736																	
		ES 2293262																	
											US 2	005-	5604	85	20051212				
PRAI		2003																	
	WO	2004	-EP6	278		W		2004	0610										

The invention relates to a preparation of 5HT4-antagonistic N-(piperidin-4-ylmethyl)-benzamide derivs. of formula I [wherein: R1 is a bivalent (un) substituted radical of formula O(CH2)1-40 or O(CH2)2-5; R2 is H or halogen; R3 is alkyl, alkoxy, or halogen; R4 is H or alkyl, and OR4 is situated at the 3- or 4-position of the piperidine ring; L is H, alkyl-(H/OH/CN/cycloalkyl), alkyl-(O/S/SO2)-(cyclo)alkyl, or alky1-C(0)-alky1, etc.]. For instance, N-(piperidin-4-ylmethy1)-benzamide derivative II (R5 = H; pIC50 = 6.97) was prepared via decarboxylation of II (R5 = t-BuO2C).

RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

Ι

- L8 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 2005:14394 CAPLUS
- DN 142:114101
 - Preparation of N-(piperidinylmethyl) benzamide derivatives as 5HT4-antagonists
- IN Bosmans, Jean-Paul Rene Marie Andre; Gijsen, Henricus Jacobus Maria; Mevellec, Laurence Anne
- PA Janssen Pharmaceutica N.V., Belg.
- PCT Int. Appl., 98 pp. SO
- CODEN: PIXXD2
- DT Patent
- T.A English

FAN.	CNT 1																
	PATENT	NO.			KIN	D	DATE			APPL	ICAT	ION :	NO.		D.	ATE	
						-											
PI	WO 2005000838				A1		20050106		WO 2004-EP6285						20040610		
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		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	ıs,	JP,	KΕ,	KG,	KP,	KR,	KZ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NA,	NI,
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
		ΤJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,
		ΑZ,	BY,	KG,	KΖ,	MD,	RU,	TJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,
		EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,
		SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,

OS CASREACT 142:114101; MARPAT 142:114101

GΙ

AB Title compds. represented by the formula I [wherein R1R2 = OCH2O, O(CH2) nOm, O(CH2) 5; n = 2-4; m = 0 or 1; R3 = H, halo or alkyl; R4 = (cyano) alkyl, alkoxy(alkyl), cyano, (alkyl) amino; R5 = H or alkyl; L = H, alkyl(cyano), alkoxyalkyl, alkylcarbamoyl, etc.; stereochem. isomers thereof, an N-oxides thereof, and pharmaceutically acceptable acid or base addition salts thereof] were prepared as 5HT4-antagonists. For example, II was given in a multi-step synthesis starting from Me 5-nitro-2,3-dihydroxybenzoate. I were tested for 5HT4 antagonistic activity with pIC50 values of around 6-9, and showed metabolic stability as well. Thus, I and their pharmaceutical compns. are useful as a medicine of 5HT4-antagonists.

RE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- 1.8 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 2005:14393 CAPLUS
- DN 142:113910

- TI Preparation of aminosulfonyl substituted 4-(aminomethyl)-piperidine benzamides as $5\mathrm{HT}4\mathrm{-antagonists}$
- IN Bosmans, Jean-Paul Rene Marie Andre; Gijsen, Henricus Jacobus Maria; Mevellec, Laurence Anne
- PA Janssen Pharmaceutica N.V., Belg.
- SO PCT Int. Appl., 53 pp.
- CODEN: PIXXD2
- DT Patent
- LA English

FAN.	PATENT NO.						KIND DATE				APPL								
PI	WO	2005	0008	37												0040	610		
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			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	
			LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,	
			NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	
			TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW	
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								GR,											
						BF,	ΒJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	ΝE,	
				TD,															
						A1 20050106				AU 2004-251823									
		2526								CA 2004-2526079 EP 2004-739781									
	EΡ	1641																	
		R:						ES,											
								RO,											HR
		2007																	
										US 2005-560300						20051212			
PRAI		2003																	
		2003																	
	WO 2004-EP6280							2004	0610										
os	MAI	RPAT	142:	1139	10														
GI																			

OR5

AB Novel compds. of formula I [X = O(CH2)nO, O(CH2)n; n = 1-5; R3 = H, halo, alkyl, alkoxy; R4 = H, halo, alkyl, alkoxy, cyanoalkyl, CN, (substituted) amino; R5 = H, alkyl; L = (substituted) aminosulfonylalkyl, alkylsulfonylaminocarbonylalkyl, etc.] are prepared which have

II

5HT4-antagonistic properties. The invention further relates to methods for preparing such compds., pharmaceutical compns. comprising said compds. as well as the use as a medicine of said compds. Thus, II was prepared, and had 5HT4 antagonism activity with pIC50 of 7.92, and was 5% metabolized

after 60 min in liver tissue. RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L8 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2008 ACS on STN
- 2000:441788 CAPLUS AN
- DN 133:74035
- ΤI Preparation of 4-(aminomethyl)piperidinebenzamides as gastrointestinal agents.
- TN Bosmans, Jean-Paul Rene Marie Andre; Meulemans, Ann Louise Gabrielle; De Cleyn, Michel Anna Jozef; Gijsen, Henricus Jacobus Maria
- PA Janssen Pharmaceutica N.V., Belg.
- PCT Int. Appl., 58 pp. SO
- CODEN: PIXXD2 DT Patent
- LA English
- F

FAN.CNT 1																					
	PA?	ENT	NO.			KIN	D	DATE		APPLICATION NO.							DATE				
PI									WO 1999-EP10064						19991214						
		W:						AZ,													
								ES,													
								KP,													
								MX,													
								TT,													
		RW:						SD,													
								GR,								SE,	BF,	ВJ,	CF,		
			CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE	Ξ,	SN,	TD,	TG						
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	BR	W 570920 CA 2355857 SR 9916491 CP 1140915 CP 1140915				A		2001	0904		BR	19	1999-16491				1	214			
	EP					A1		2001	1010		EP 1999-967956						1	214			
	EP																				
		R:						ES,	FR,	GB,	GE	₹,	IT,	LI,	LU,	NL,	SE,	MC,	PT,		
			IE,	SI,	LT,	LV,	FI,	RO													
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					A2		2002	0729		HU	20)0I-	4838			1	9991	214			
	HU	2001	0048	38		A3		2003	0528			~		225					02.4		
	TD	2001	0033	27		A.		2002	1000		TD	20	JU1	333	2.2		1	9991	214		
	NE	E120	71	3 /		2		2002	1100		NE	20	000-	5120	71		1	0001	214		
	NZI NII	7702	07			D2		2002	0210		NZ.	13	299-	2120	/ I		1	0001	214		
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	AT	2979	17			т		2005	0715		AT	10	999-	9679	56		1	9991	214		
	PT	1140	915			Ť		2005	1130		PT	10	999-	9679	56		1	9991	214		
	ES	2245	131			тз		2005	1216		ES	19	999-	9679	56		1	9991	214		
	SK	2858	29			B6		2007	0906		SK	20	001-	859			1	9991	214		
	PI.	1974	09			B1		2008	0331		PI.	10	999-	3484	17		1	9991	214		
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	BG	6495	3			A B1		2006	1031												
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	HR 2001000445					A1		2002	0630		HR	20	001-	445			2	0010	614		
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	ZA	2001	0051	35		A		2002	0621		ZA	20	001-	5135			2	0010	621		

HK 1039114 US 20030181456 US 7205410 PRAI EP 1998-204411 WO 1999-EP10064	A1 A1 B2 A W	20050819 20030925 20070417 19981222 19991214	HK 2002-100161 US 2003-353307	20020129 20030129
US 2001-857905 OS MARPAT 133:74035	A3	20010608		
OD MAREAI 133.74033				

AB Title compds. [I; RIR2 = (substituted) OCH2CH2, OCH2CH2, OCH2CH2O, etc.; R3 = H, halo; R4, R5 = H, alkyl; L = cycloalkyl, oxocycloalkyl, alkenyl, etc.], were prepared Thus, trans-N-[1-(3-aminopropyl)-3-hydroxy-4-piperidinyl]methyl-7-chloro-2,3-dihydro-1,4-benzodioxin-5-carboxamide (preparation given). Was stirred with 2-chloro-3-methylpyrazine and CaO at 120° to give 16% trans-7-chloro-2,3-dihydro-N-[(3-hydroxy-1-[3-[(3-methyl-2-pyrazinyl)amino]propyl1-4-piperidinyl]methyl-1,4-benzodioxin-5-carboxamide. This antagonized 5HT4 in rat esophageal tunica muscularis mucosae with pA2 = 10.55.

RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT